

Europäisches Patentamt

European Patent Office

Office européen des brevets



EP 1 092 551 A2

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication:

18.04.2001 Bulletin 2001/16

(51) Int Cl.7: B41M 5/30

(11)

(21) Application number: 00309043.8

(22) Date of filing: 13.10.2000

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 15.10.1999 JP 29394799

13.12.1999 JP 35315999

(71) Applicant: NIPPON PAPER INDUSTRIES CO., LTD. Kita-ku, Tokyo 114-0002 (JP)

(72) Inventors:

Hamada, Kaoru, c/o Res. Lab.,
 Product Development
 1-chome, Shinjuku-ku, Tokyo 161-0034 (JP)

Tadakazu, Fukuchi, Res. Lab.,
 Product Developm.
 1-chome, Shinjuku-ku, Tokyo 161-0034 (JP)

Midorikawa, Yoshimi c/o Res. Lab.,
 Product Develop
 1-chome, Shinjuku-ku, Tokyo 161.0034 (JP)

• Kimura, Yoshihide, c/o Res. Lab.,

Product Develop. 1-chome, Shinjuku-ku, Tokyo 161-0034 (JP)

(74) Representative: McCluskie, Gail Wilson
J.A. Kemp & Co.,
14 South Square,
Gray's Inn
London WC1R 5LX (GB)

(54) Multi chromatic thermally sensitive recording medium

(57) A multi chromatic thermally sensitive recording medium comprising, a higher temperature color developing layer containing dye precursor and an organic color developer that reacts with said dye precursor and develops color by heating, and at least one lower temperature color developing layer that develops different color from that of said higher temperature color developing layer and develops color by lower temperature than the color developing temperature of said higher temperature color developing layer laminated in order on a substrate, wherein an organic color developer contained in said higher temperature color developing layer contains said diphenylsulfone bridgeable type compound represented by general formula (1) and/or said 3-{[(phenylamine)carbonyl] amino}benzenesulfoneamide compound represented by general formula (2).

$$(R_1)_m \qquad (R_2)_n \qquad (R_3)_p \qquad (R_4)_q \qquad (R_5)_r \qquad (R_6)_t \qquad (R_6)_t \qquad (R_7)_p \qquad (R_8)_p \qquad (R_8$$

wherein. X and Y can be different and indicates a saturate or an unsaturated liner or grafted hydrocarbon group of carbon number 1 to 12 which can possess an ether bond, or

$$-R$$
 (a)

or.

$$-CH2-C-CH2-C (b)$$

wherein R indicates a methylene group or an ethylene group, T indicates a hydrogen atom or an alkyl group of carbon number 1 to 4.

and R_1 to R_6 independently indicate a halogen atom, an alkyl group of carbon number 1 to 6 or an alkenyl group, further m, n, p, q, r, t indicate an integer number of 0 to 4 and when are bigger than 2, R_1 to R_6 can be different and a is an integer of 0 to 10.

Description

10

15

25

35

45

50

55

[0001] The present invention relates to a multi chromatic thermally sensitive recording medium, more in detail relates to a multi chromatic thermally sensitive recording medium having plural color developing layer which develops different color tones by different heating conditions of thermal head.

[0002] In general, a thermally sensitive recording media having a thermally sensitive color developing layer mainly composed of colorless or pale colored electron donning type dye precursor (hereinafter shortened to dye precursor) and color developer that develops color by reacting with dye precursor when heated, is disclosed in Japanese Patent Publication 45-14035 and is widely used in a commercial scale. As a method for recording of this thermally sensitive recording medium, usually, a thermal printer in which a thermal head is installed is used. This kind of thermal recording method is superior to a conventional recording method from the view point of noiseless at recording process, does not need a developing and fixing processes, maintenance free, equipment is relatively cheap and compact and the obtained image is very clear. Therefore, this method is widely applied in the field of a facsimile or a computer, various kinds of measuring instrument and for a labeling machine along with the growth of an information industry. Along with the expanding of the usage, the required qualities to a thermally sensitive recording medium are becoming more multiplex, for instance, a higher sensitivity, a stabilization of image and a multi coloration of image can be mentioned. Especially, the multi coloration of recorded image have a merit that the letters or patterns to be emphasize can be recorded by a different color tone from the other part.

[0003] As a multi chromatic thermally sensitive recording medium, for example, a method to prepare plural color developing layers that develop different colors on a substrate and to form a recording image using variations of heating temperature or thermal energy can be mentioned. The structural feature of said thermally sensitive recording medium is characterized to laminate the layers in order from a higher temperature color developing layer to a lower temperature color developing layers that develops color by lower temperature or lower thermal energy than said higher temperature color developing layer. These multi chromatic thermally sensitive recording media can be mainly divided to two different types, that is, a color discharging type and a color adding type.

[0004] The color developing mechanism of the color discharging type is disclosed in e.g. Japanese Patent Publication 50-17865, Japanese Patent Publication 57-14320 or Japanese Patent Laid Open Publication 2-80287. That is, only a lower temperature color developing layer develops color by lower temperature heating, and when heated by higher temperature, the color developed by lower temperature is discharged by the action of color discharging agent having color discharging effect, and only a higher color developing layer develops color. This method has a merit that the desired color tone can be selected voluntarily, however, it is necessary to add large amount of color discharging agent to perform a sufficient color discharging effect. The color discharging type multi chromatic thermally sensitive recording medium have problems, that is, the deterioration of recorded image preservability caused by added large amount of color discharging agent or the deterioration of recording sensitivity caused by a large consumption of thermal energy necessary for melting the achromatic agent.

[0005] Meanwhile, as the example of a color adding type thermally sensitive recording medium, the methods disclosed in Japanese Patent Publication 49-27708, Japanese Patent Publication 51-19989 or Japanese Patent Laid Open Publication 51-146239 can be mentioned. That is, the methods to obtain distinguishable two color images by adding different thermal energy to a double layered color developing layer which develops different two colors are disclosed. In the methods disclosed above, a lower temperature developing layer located at the upper position develops color by lower heating temperature, and a higher temperature developing layer located at the bottom position develops color by higher heating temperature, thus both two color developing layers develop colors. And, since in the obtained image these two colors of said two layers are mixed, it is substantially necessary to make the color of the layer located at the bottom position black. Further, since the black color image of bottom layer is obtained by mixing with the color of an upper layer, the color of an upper layer is fogged over the black color image and the difference between said two developed color tones is not so clear. Furthermore, there are problems that when the developed color is exposed to the high temperature atmosphere, for example on a dash board of car, the higher temperature developing layer develops color and a color tone developed by lower temperature becomes fogging or becomes black. In Japanese Patent Laid Open Publication 4-329186, there is a description that the color cloudy problem caused by color developing of a higher temperature color developing layer at the lower temperature heating can be solved by containing specific color indicator in a higher temperature color developing layer. However, this method is not sufficient at the clearness of color tone developed at lower temperature in high temperature atmosphere and the stability of ground color. Still further, recently, the case to use a thermally sensitive recording paper as a receipt becomes more popular, in which case the preservability of recorded image becomes necessary. However, a thermally sensitive recording paper having sufficient image preservability is not yet developed up to the present time.

[0006] The object of this invention is to provide a multi chromatic thermally sensitive recording medium that can obtain clear color developed images both at a lower temperature color developing layer and a higher color developing layer, further the obtained color image developed at lower temperature does not become fogging by the color tone of

higher temperature color developing layer at the high temperature atmosphere, furthermore the image preservability is excellent.

[0007] The present invention is a multi chromatic thermally sensitive recording medium comprising, a higher temperature color developing layer containing dye precursor and an organic color developer that reacts with said dye precursor and develops color by heating, and at least one lower temperature color developing layer that develops different color from that of said higher temperature color developing layer and develops color by lower temperature than the color developing temperature of said higher temperature color developing layer laminated in order on a substrate, wherein an organic color developer contained in said higher temperature color developing layer contains at least one compound selected from the group consisted of diphenylsulfone bridgeable type compound represented by general formula (1), which is disclosed in WO97/16420, and 3-{[(phenylamine)carbonyl] amino}benzenesulfoneamide compound represented by general formula (2), which is disclosed in Japanese Patent Laid Open Publication 8-59603.

$$(R_1)_m \qquad (R_2)_n \qquad (R_3)_p \qquad (R_4)_q \qquad (R_5)_r \qquad (R_6)_l$$

$$(R_1)_m \qquad (R_2)_n \qquad (R_3)_p \qquad (R_4)_q \qquad (R_5)_r \qquad (R_6)_l \qquad (R_6$$

wherein. X and Y can be different and indicates a saturate or an unsaturated liner or grafted hydrocarbon group of carbon number 1 to 12 which can possess an ether bond, or

or

10

25

35

45

50

55

$$-CH_2 - C - CH_2 - CH_2 - (b)$$

wherein R indicates a methylene group or an ethylene group, T indicates a hydrogen atom or an alkyl group of carbon number 1 to 4.

and R_1 to R_6 independently indicate a halogen atom, an alkyl group of carbon number 1 to 6 or an alkenyl group, further m, n, p, q, r, t indicate an integer number of 0 to 4 and when are bigger than 2, R_1 to R_6 can be different and a is an integer of 0 to 10.

[0008] Further, in the present invention, the lower temperature color developing layer which develops different color from that of the higher temperature color developing layer and develops color by lower temperature than that of the

color developing temperature of said higher temperature color developing layer is not restricted to only one layer, and laminated from higher ones to lower ones in order on a substrate, thus the multi chromatic thermally sensitive recording medium can be obtained.

[0009] The multi chromatic thermally sensitive recording medium which develops more than 2 colors can be easily affected by variations of temperature environment, because of it's color developing mechanism which used difference of heating temperature or thermal energy. In a case of the color tone developed by lower temperature formed by lower temperature heating (lower energy printing), for the purpose to maintain the cleamess of color tone in higher level, it is important to control the effect of color developing of the higher temperature color developing layer to the lowest level. As the countermeasure to prevent the partial color developing of the higher temperature color developing layer at the lower heating temperature, the raising of the melting point of the color developer used in the higher temperature color developing layer is concerned to be effective.

[0010] However, even if the color developer having higher melting point is used, at the lamination process of a higher color developing layer and a lower color developing layer, both layers are partially mixed at the boarder, and generates melting point dropping phenomenon. Therefore, compared with the cases that a higher temperature color developing layer or a lower temperature color developing layer is independently prepared on the substrate, the stability of ground color to a heat is remarkably deteriorated and also the whiteness becomes bad. For example, in cases when 2,4'-dihydroxydiphenylsulfone or 4,4'-dihydroxy diphenylsulfone which are the conventional well-known color developer having high melting point, since the color developing of ground color is generated at the atmosphere of 60°C around temperature, the stability of the ground color is not so sufficient.

[0011] In the meanwhile, the inventors of this invention have found that by containing at least one kind of compound selected from the group consisted of diphenylsulfone type bridgeable compound represented by general formula (1) or 3-{[(phenylamine)carbonyl]amino}benzenesulfoneamide compound represented by general formula (2) in a higher temperature color developing layer, the ground color development at high temperature atmosphere is prevented, the fogging to the lower temperature developed color image by the higher temperature color developing layer is improved, and further the preservability of the image developed at higher temperature is remarkably improved, and accomplished the present invention. The reason why the excellent function and effect are obtained by the present invention is not made clear. By the way, diphenylsulfone type bridgeable type compound represented by general formula (1) and 3-{[(phenylamine)carbonyl]amino}benzenesulfone -amide compound represented by general formula (2) are especially superior in the stability to heat compared with the conventional phenol type color developer. Therefore, when they are contained in a higher temperature color developing layer, the fogging by the higher temperature color developing layer to the image developed by lower temperature becomes small, further do not develop color even if they are placed in high temperature atmosphere. And it is concerned that this is the reason why the multi chromatic thermally sensitive recording medium of this invention can maintain the color tone of the lower temperature color developed image clearly. Further, when the diphenylsulfone type bridgeable type compound represented by general formula (1) is use, the stability of obtained image to the solvent such as oil or plasticiser are remarkably higher compared with that of conventional phenol type color developer. The reason why is not clear yet, however, the presumed reasons are mentioned below. First, the bonding strength with leuco dye is high because it has plural numbers of sulfonic group in the molecular, second, the solubility of a kind of complex formed when it reacts with leuco dye to these solvents is low.

[0012] In the multi chromatic thermally sensitive recording medium of this invention, the color tone of each color developing layer are adjusted by the selection of leuco dye which is a dye precursor. And, for example, in a case of two colored thermally sensitive recording medium, it is desired to select a mono tone color (e.g. blue, red or yellow) dye whose absorption peak after color developed is single for a lower temperature color developing layer and to select a multi tone color (e.g. green or black) dye whose absorption peak after developed by mixed color tone is plural for a higher temperature color developing layer.

[0013] In a case to use a black type leuco dye to a higher temperature color developing layer of this invention, especially

3-diethylamino-7-(o-chloroanilino)fluoran,

10

15

25

35

45

50

55

- 3-diethylamino-6-methyl-7-anilinofluoran,
- 3-dibutylamino-6-methyl-7-anilinofluoran,
- 3-dibutylamino-7-(o-chloroanilino)fluoran,
- 3-di-n-pentylamino-6-methyl-7-anilinofluoran,
- 3-(N-ethyl-p-toluidino)-6-methyl-7-anilinofluoran and
- 3-(N-ethyl-N-isoamylamino)-6-methyl-7-anilinofluoran

are desirably used because when used together with diphenylsulfone type bridgeable type compound represented by general formula (1) or 3-{[(phenylamine)carbonyl]amino}benzenesulfoneamide compound represented by general formula (2) do not develop color at lower temperature heating and do not cause the problem of fogging to the image

developed by lower temperature, further, very clear black colored image can be obtained at high temperature heating. Especially, among these compounds following compounds, that is.

- 3-diethylamino-7-(o-chloroanilino)fluoran.
- 3-dibutylamino-7-(o-chloroanilino)fluoran and
- 3-(N-ethyl-p-toluidino)-6-methyl-7-anilinofluoran

are desirably used, because a thermally sensitive recording medium whose ground color developing at the preservation in high temperature condition becomes very small can be obtained.

[0014] Besides above mentioned compounds, as the leuco dye used in the present invention, the conventional well known dyes in the field of a pressure sensitive type or a thermally sensitive recording paper can be used. Desirably, triphenyl methane type compound, fluoran type compound, fluorene type compound and divinyl type compound can be used, however, not intended to be limited to them. Typical examples of basic colorless dye are mentioned below. Further, these dye precursors can be used alone or together with.

<Triphenylmethane type leuco dyes>

[0015]

10

15

20

35

50

55

- 3,3-bis(p-dimethylaminophenyl)-6-dimethylaminophthalide (another name is Crystal Violet Lactone)
- 3,3-bis(p-dimethylaminophenyl)phthalide [another name is Malachite Green Lactone]

<Fluoran type leuco dyes>

25 [0016]

- 3-diethylamino-6-methylfluoran
- 3-diethylamino-6-methyl-7-anilinofluoran
- 3-diethylamino-6-methyl-7-(o,p-dimethylanilino)fluoran
- 30 3-diethylamino-6-methyl-7-chlorofluoran
 - 3-diethylamino-6-methyl-7-(m-trifluoromethylanilino)fluoran
 - 3-diethylamino-6-methyl-7-(m-methylanilino)fluoran
 - 3-diethylamino-6-methyl-7-(o-chloroanilino)fluoran
 - 3-diethylamino-6-methyl-7-(p-chloroanilino)fluoran
 - 3-diethylamino-6-methyl-7-(o-fluoroanilino)fluoran
 - 3-diethylamino-6-methyl-7-n-octylanilinofluoran
 - 3-diethylamino-6-methyl-7-n-octylaminofluoran
 - 3-diethylamino-6-methyl-7-benzylanilinofluoran
 - 3-diethylamino-6-methyl-7-dibenzylanilinofluoran
- 40 3-diethylamino-6-chloro-7-methylfluoran
 - 3-diethylamino-6-chloro-7-anilinofluoran
 - 3-diethylamino-6-chloro-7-p-methylanilinofluoran
 - 3-diethylamino-6-ethoxyethyl-7-anilinofluoran
 - 3-diethylamino-7-methylfluoran
- 45 3-diethylamino-7-dibenzylaminofluoran
 - 3-diethylamino-6,8-dimethylfluoran
 - 3-diethylamino-7-chlorofluoran
 - 3-diethylamino-7-(m-trifluoromethylanilino)fluoran
 - 3-diethylamino-7-(o-chloroanilino)fluoran
 - 3-diethylamino-7-(p-chloroanilino)fluoran
 - 3-diethylamino-7-(o-fluoroanilino)fluoran
 - 3-diethylamino-benzo[a]fluoran
 - 3-diethylamino-benzo[c]fluoran
 - 3-dibutylamino-6-methyl-fluoran
 - 3-dibutylamino-6-methyl-7-anilinofluoran
 - 3-dibutylamino-6-methyl-7-(o.p-dimethylanilino)fluoran
 - 3-dibutylamino-6-methyl-7-(o-chloroanilino)fluoran
 - 3-dibutylamino-6-methyl-7-(p-chloroanilino)fluoran

	3-dibutylamino-6-methyl-7-(o-fluoroanilino)fluoran
	3-dibutylamino-6-methyl-7-(m-trifluoromethylanilino)fluoran
	3-dibutylamino-6-methyl-7-chlorofluoran
	3-dibutylamino-6-methyl-7-bromofluoran
5	3-dibutylamino-6-ethoxyethyl-7-anilinofluoran
	3-dibutylamino-6-chloro-7-anilinofluoran
	3-dibutylamino-6-methyl-7-p-methylanilinofluoran
	3-dibutylamino-7-(o-chloroanilino)fluoran
	3-dibutylamino-7-(o-fluoroanilino)fluoran
10	3-di-n-pentylamino-6-methyl-7-anilinofluoran
	3-di-n-pentylamino-6-methyl-7-(p-chloroanilino)fluoran
	3-di-n-pentylamino-7-(m-trifluoromethylaniliono)fluoran
	3-di-n-pentylamino-6-chloro-7-anilinofluoran
	3-di-n-pentylamino-7-(p-chloroanilino)fluoran
15	3-pyrrolidino-6-methyl-7-anilinofluoran
	3-piperidino-6-methyl-7-anilinofluoran
	3-(N-methyl-N-propylamino)-6-methyl-7-anilinofluoran
	3-(N-methyl-N-cyclohexylamino)-6-methyl-7-anilinofluoran
	3-(N-ethyl-N-cyclohexylamino)-6-methyl-7-anilinofluoran
20	3-(N-ethyl-N-xylamino)-6-methyl-7-(p-chloroanilino)fluoran
	3-(N-ethyl-p-toluidino)-6-methyl-7-anilinofluoran
	3-(N-ethyl-N-isoamylamino)-6-methyl-7-anilinofluoran
	3-(N-ethyl-N-isoamylamino)-6-chloro-7-anilinofluoran
	3-(N-ethyl-N-tetrahydrofurfurylamino)-6-methyl-7-anilinofluoran
25	3-(N-ethyl-N-isobutylamino)-6-methyl-7-anilinofluoran
	3-(N-p-tolyl-N-ethylamino)-7-(N-phenylmethyl-N-methyl)fluoran
	3-(N-ethyl-N-ethoxypropylamino)-6-methyl-7-anilinofluoran
	3-N-ethyl-N-isoamylamino-benzo[a]fluoran
	3-N-ethyl-N-p-methylphenylamino-7-methylfluoran
30	3-cyclohexylamino-6-chlorofluoran
	2-(4-oxahexyl)-3-dimethylamino-6-methyl-7-anilinofluoran
	2-(4-oxahexyl)-3-diethylamino-6-methyl-7-anilinofluoran
	2-(4-oxahexyl)-3-dipropylamino-6-methyl-7-anilinofluoran
	2-methyl-6-p-(p-dimethylaminophenyl)aminoanilinofluoran
35	2-methoxy-6-p-(p-dimethylaminophenyl)aminoanilinofluoran
	2-chloro-3-methyl-6-p-(p-phenylaminophenyl)aminoanilinofluorar
	2-chloro-6-p-(p-dimethylaminophenyl)aminoanilinofluoran
	2-nitro-6-p-(p-diethylaminophenyl)aminoanilinofluoran
	2-amino-6-p-(p-diethylaminophenyl) aminoanilinofluoran
40	2-diethylamino-6-p-(p-diethylaminophenyl)aminoanilinofluoran
	2-phenyl-6-metyl-6-p-(p-phenylaminophenyl)aminoanilinofluoran
	2-benzyl-6-p-(p-phenylaminophenyl)aminoanilinofluoran
	2-hydroxy-6-p-(p-phenylaminophenyl)aminoanilinofluoran
	3-methyl-6-p-(p-dimethylaminophenyl)aminoanilinofluoran
45	3-diethylamino-6-p-(p-diethylaminophenyl)aminoanilinofluoran
	3-diethylamino-6-p-(p-dibutylaminophenyl)aminoanilinofluoran
	2,4-dimethyl-6-[(4-dimethylamino)anilino]-fluoran
	<fluorene dyes="" leuco="" type=""></fluorene>
50	

[0017]

3,6,6'-tris(dimethylamino)spiro[fluorene-9,3'-phthalide] 3,6,6'-tris(diethylamino)spiro[fluorene-9,3'-phthalide]

<Divinyl type leuco dyes>

[0018]

- $3.3-bis-[2-(p-dimethylaminophenyl)-2-(p-methoxyphenyl)\\ ethenyl]-4,5,6,7-tetra\ bromophthalide$
- 3.3-bis-[2-(p-dimethylaminophenyl)-2-(p-methoxyphenyl)ethenyl]-4,5,6,7-tetra chlorophthalide
- 3.3-bis-(1.1-bis(4-pyrrolidinophenyl)ethylen-2-yll-4,5,6,7-tetrabromophthalide
- 3.3-bis-[1-(4-methoxyphenyl)-1-(4-pyrrolidinophenyl)ethylene-2-yl]-4,5,6,7-tetra chlorophthalide

10 <Others>

15

20

25

30

40

45

50

[0019]

- 3-(4-diethylamino-2-ethoxyphenyl)-3-(1-ethyl-2-methylindol-3-yl)-4-aza phthalide
- 3-(4-diethylamino-2-ethoxyphenyl)-3-(1-octyl-2-methylindol-3-yl)-4-aza phthalide
- 3-(4-cyclohexylethylamino-2-methoxyphenyl)-3-(1-ethyl-2-methylindol-3-yl)-4-azaphthalide
- 3,3-bis(1-ethyl-2-methylindol-3-yl)phthalide
- 3.3-bis(1-n-butyl-2-methylindol-3-yl)phtalide
- 3.3-bis(1-n-octyl-2-methylindol-3-yl)phtalide
- 3,6-bis(dietylamino)fluoran-y-anilinolactam
- 3,6-bis(diethylamino)fluoran-y-(3'-nitro)anilinolactam
- 3,6-bis(diethylamino)fluoran-y-(4'-nitro)anilinolactam
- 1,1-bis-[2',2',2",2"-tetrakis-(p-dimethylaminophenyl)-ethenyl]-2,2-dinitrilethane
- 1,1-bis-[2',2',2",2"-tetrakis-(p-dimethylaminophenyl)-ethenyl]-2-β-naphthoyl ethane
- 1,1-bis-[2',2',2",2"-tetrakis-(p-dimethylaminophenyl)-ethenyl]-2,2-diacetylethane bis-[2,2,2',2'-tetrakis-(p-dimethylaminophenyl)-ethenyl]-methylmalonicacid dimethylester.

[0020] In the present invention, as a sensitizer contained in a high temperature color developing layer, at least one kind of diphenylsuofone bridgeable type compound represented by general formula (1) is used. The diphenylsuofone bridgeable type compound represented by general formula (1) is a compound disclosed in International Publication WO97/16420 and Japanese Patent Laid Open Publication 10-29969.

[0021] In general formula (1), as the concrete examples of groups indicated by X and Y, following compounds can be mentioned. Namely, methylene group, ethylene group, trimethylene group, tetramethylene group, pentamethylene group, hexamethylene group, hexamethylene group, hexamethylene group, hexamethylene group, decamethylene group, undecamethylene group, dodecamethylene group, methylmethylene group, dimethylene group, methylene group, methylene group, methylene group, tethylene group, 1,2-dimethylethylene group, 1-methyltrimethylene group, 1-methyltrimethylene group, 1-methyltetramethylene group, vinylene group, propenylene group, 2-butenylene group, ethylene group, 2-butynylene group, 1-vinylethylene group, ethyleneoxyethylene group, tetramethyleneoxyethylene group, 1,3-dioxane-5, 5-bismethylene group, 1,2-xylyl group, 1,3-xylyl group, 1,4-xylyl group, 2-hydroxytrimethylene group, 2-hydroxy-2-methyltrimethylene group, and 2-hydroxy-2-ethyltrimethylene group can be mentioned.

10 mg 10 mg

[0022] Alkyl group or alkenyl group of R₁ to R₆ is an alkyl group of C₁ to C₆ or an alkenyl group of C₁ to C₆, and as concrete examples, methyl group, ethyl group, n-propyl group, isopropyl group, n-butyl group, sec-butyl group, tert-butyl group, isopentyl group, neopentyl group, tert-pentyl group, n-hexyl group, isohexyl group, 1-metylpentyl group, 2-methylpentyl group, vinyl group, allyl group, isopropenyl group, 1-propenyl group, 2-butenyl group, 3-butenyl group, 1,3-butandienyl group and 2-methyl-2-propenyl group can be mentioned. And as a halogen atom, chloride, bromine, fluorine or iodine are mentioned.

[0023] In this invention, as a diphenylsulfone bridgeable type compound represented by general formula (1), several kinds of compounds whose substitution group and/or "a" number are different can be used by mixing together with, and the mixing ratio can be voluntarily selected. The mixing method is not restricted, however, substantially, mixing by powder state, mixing in aqueous solution or mixing by to producing plural kinds of diphenylsulfone bridgeable type compounds simultaneously by using specific producing method can be mentioned.

[0024] As the concrete examples of a compound represented by general formula (1), the following compounds can be mentioned.

(1-1)

```
4.4'-bis[4-[4-(4-hydroxyphenylsulfonyl)phenoxy]-2-trans-butenyloxy]diphenyl sulfone
                  4.4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy-4-butyloxy]diphenylsulfone
                  (1-3)
                  4.4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy-3-propyloxy]diphenylsulfone
                  4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy-2-ethyloxy]diphenylsulfone
                  (1-5)
                  4-[4-(4-hydroxyphenylsulfonyl)phenoxy-4-butyloxy]-4'-[4-(4-hydroxyphenyl sulfonyl)phenoxy-3-propyloxy]diphe-
10
                  nylsulfone
                  (1-6)
                  4-[4-(4-hydroxyph enylsulfonyl) phenoxy-4-butyloxy]-4'-[4-(4-hydroxyphenyl sulfonyl) phenoxy-2-ethyloxy]diphe-
                  nylsulfone
                  4-[4-(4-hydroxyphenylsulfonyl) phenoxy-3-propyloxy]-4'-[4- (4-hydroxyphenyl sulfonyl) phenoxy-2-ethyloxy]diph
15
                  enylsulfone
                  (1-8)
                  4-4'-bis[4-(4-hydroxyphenylsulfonyl) phenoxy-5-pentyloxy]diphenylsulfone
                  4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy-6-hexyloxy]diphenylsulfone
20
                  4-[4-[4-hydroxyphenylsulfonyl]phenoxy]-2-trans-butenyloxy]-4'-[4-(4-hydroxy phenylsulfonyl) phenoxy-4-buty-
                  loxy]diphenylsulfone
                   (1-11)
                  4-[4-(4-hydroxyphenylsulfonyl) phenoxy-2-trans-butenyloxy]-4'- [4- (4-hydroxy phenylsulfonyl) phenoxy-3-propy-
25
                  loxy]diphenylsulfone
                   (1-12)
                   4-[4-[4-(4-hydroxyphenylsulfonyl) phenoxy]-2-trans-butenyloxy]-4'-[4-(4-hydroxy phenylsulfonyl) phenoxy-2-ethy-
                   loxy]diphenylsulfone
30
                   (1-13)
                   1,4-bis[4-[4-(4-hydroxyphenylsulfonyl)phenoxy-2-trans-butenyloxy]phenyl sulfonyl]phenoxy]-cis-2-butene
                   (1-14)
                   1,4-bis[4[4-[4-(4-hydroxyphenylsulfonyl) phenoxy-2-trans-butenyloxy]phenyl sulfonyl]phenoxy]-trans-2-butene
                   (1-15)
                   4;4'-bis[4-[4-(2-hydroxyphenylsulfonyl) phenoxy]butyloxy]diphenylsulfone
                                                                                                                                                                                                                          en y a
35
                   4,4'-bis[4-[2-(4-hydroxyphenylsulfonyl) phenoxy]butyloxy]diphenylsulfone
                   (1-17)
                   4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy-2-ethylenoxyethoxy]diphenyl sulfone
40
                   4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenyl-1,4-phenylenebismethyleneoxy] diphenylsulfone,
                   4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenyl-1,3-phenylenebismethyleneoxy] diphenylsulfone
                   4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenyl-1,2-phenylenebismethyleneoxy] diphenylsulfone
45
                   2,2'-bis[4-[4-[4-(4-hydroxyphenylsulfonyl) phenoxy-2-ethyleneoxyethoxy]phenyl sulfonyl] phenoxy] diethyl ether
                   a, a '-bis[4-[4-(4-hydroxyphenylsulfonyl)phenyl-1.4-phenylenebismethylene oxy] phenylsulfonyl]phenoxy]-p-
50
                   \alpha,\alpha'-bis[4-[4-(4-hydroxyphenylsulfonyl)phenyl-1,3-phenylenebismethylene oxy]phenylsulfonyl]phenoxy]-m-xy-
                   lene
                   (1-24)
                   \alpha,\alpha'-bis[4-[4-(4-hydroxyphenylsulfonyl]phenyl-1,2-phenylenebismethylene \\ oxy]phenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenoxy]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenylsulfonyl]-0-xy-bis[4-[4-(4-hydroxyphenylsulfonyl]phenylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsulfonylsu
55
                    (1-25)
```

2.4'-bis[2-(4-hydroxyphenylsulfonyl)phenoxy-2-ethyleneoxyethoxy] diphenylsulfone

(1-26)

2.4'-bis[4-(2-hydroxyphenylsulfonyl)phenoxy-2-ethyleneoxyethoxy] diphenylsulfone

(1-27)

4.4'-bis[3.5-dimethyl-4-(3.5-dimethyl-4-hydroxyphenylsulfonyl) phenoxy-2-ethylene oxyethoxy]diphenylsulfone (1-28)

4.4'-bis[3-allyl-4-(3-allyl-4-hydroxyphenylsulfonyl)phenoxy-2-ethyleneoxyethoxy] diphenylsulfone

(1-29)

4,4'-bis[3,5-dimethyl-4-(3.5-dimethyl-4-hydroxyphenylsulfonyl) phenyl-1,4-phenylenebismethyleneoxy]diphenyl-sulfone

(1-30)

10

15

20

25

30

50

55

4,4'-bis[3,5-dimethyl-4-(3.5-dimethyl-4-hydroxyphenylsulfonyl)phenyl-1,3-phenylenebismethyleneoxy]diphenyl-sulfone

(1-31)

4,4'-bis[3,5-dimethyl-4-(3.5-dimethyl-4-hydroxyphenylsulfonyl) phenyl-1,2-phenylenebismethyleneoxy]diphenyl-sulfone

(1-32)

4,4'-bis[3-allyl-4-(3-allyl-4-hydroxyphenylsulfonyl)1,4-phenylenebismethylene oxy]diphenylsulfone

(1-33)

4,4'-bis[3-allyl-4-(3-allyl-4-hydroxyphenylsulfonyl)1,3-phenylenebismethylene oxy]diphenylsulfone

(1-34)

4,4'-bis[3-allyl-4-(3-allyl-4-hydroxyphenylsulfonyl)1,2-phenylenebismethylene oxy]diphenylsulfone

(1-35)

4.4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy-2-hydroxypropyloxy]diphenyl sulfone

(1-36)

1,3-bis[4-[4-(4-hydroxyphenylsulfonyl)phenoxy-2-hydroxypropyloxy]phenyl sulfonyl]phenoxy]-2-hydroxypropane.

[0025] In a case to use by mixing several kinds of diphenylsulfone bridgeable type compound represented by general formula (1), the most desirable composition is the composition containing more than two kinds of compound represented by general formula (3) only whose "a" values are different. The preparation methods of these kind of compounds are not so complicated, and by altering the reaction ratio of materials the compounds whose "a" values are different can be synthesized by voluntary containing ratio at one time.

$$(R_1)_{m} \qquad (R_1)_{m} \qquad (R_1$$

In this formula, X. Y. R₁, m and a is same as to mentioned above.

[0026] Among the compounds represented by general formula 3, especially the compound of a=0 is a compound disclosed in W093/06074 and WO95/33714, and as the concrete example,

1,3-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]-2-hydroxypropane,

1,1-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]methane,

1,2-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]ethane,

1,3-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]propane.

1,4-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]butane.

1,5-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]pentane.

1.6-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]hexane.

a, α'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]-p-xylene,

a, α'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]-m-xylene,

a. α' -bis[4-(4-hydroxyphenylsulfonyl)phenoxy}-o-xylene.

2.2'-bis[4-(4-hydroxyphrnylsulfonyl)phenoxy]diethyl ether,

4.4'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]dibuthyl ether.

1.2-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]ethylene and

1.4-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]-2-butene can be mentioned.

[0027] The compound represented by general formula (1), can be obtained by the method described in International Patent Laid Open Publication WO97/16420 which reacts 4,4'-dihydroxyphenylsulfone derivatives or 2,4'-dihydroxyphenyl sulfone derivatives under the presence of basic compound. The color developer used in this invention contains one or more kinds of diphenylsulfone bridgeable type compound prepared by above mentioned method, and especially the compounds obtained by following synthetic examples are desirably used.

Synthetic Example 1

[0028] 16.0g (0.4 mole) of sodium hydroxide is added to 21.2g of water and dissolved, and then 50.0g (0.2 mole) of 4.4'-dihydroxydiphenylsulfone (hereinafter shortened to BPS) is added. Then, 14.3g (0.10 mole) of bis(2-chloroethyl) ether is added at 105°C, and reacted for 5 hours at 110-115°C. After the reaction is over, 375ml of water is added to the reacted solution, stirred for 1 hour at 90°C. Then cooled down to the room temperature, neutralized by 20% sulfuric acid. The crystallized solid is filtrated, and 39.3g of white crystalline is obtained. The yield to bis(2-chloroethyl)ether is 88°o. The obtained component is analyzed by high performance liquid chromatography and identified as follows. As the column, Mightysil RP-18 (product of Kanto Chemical Co., Ltd.) is used, and moving bed is CH₃CN:H₂O: 1°oH₃PO₄=700:300:5, and UV wave length is 260nm.

$$SO_2$$
 SO_2 SO_2 SO_2 SO_2 SO_2 SO_2 SO_2

X=CH2CH2OCH2CH2

30

35

40

50

55

10

20

25

```
a=0: retention time 1.9 minutes: area % 32.9
a=1: retention time 2.3 minutes: area % 21.7
a=2: retention time 2.7 minutes: area % 12.8
a=3: retention time 3.4 minutes: area % 8.8
a=4: retention time 4.2 minutes: area % 5.8
a=5: retention time 5.4 minutes: area % 3.5
a=6: retention time 7.0 minutes: area % 2.2
a=7: retention time 9.0 minutes: area % 1.7
a=8: retention time 11.8 minutes: area % 1.3
a=9: retention time 15.4 minutes: area % 1.3
```

Synthetic Examples 2-4

[0029] The molar ratio of BPS and bis(2-chloroethyl)ether of Synthetic Examplel is changed to 1.5:1, 2.5:1, 3.0:1, and following composition can be obtained. In a case of 1.5:1,

a=0;area%:20.8, a=1 ;area%:33.0, a=2;area%:14.2, a=3;area%:7.9, a=4; area%:3.9 In a case of 2.5:1,

a=0;area%:49.6, a=1;area%:25.9, a=2;area%:11.4, a=;area%:5.3, a=4;area%:3.4 In a case of 3.0:1,

a=0;area%:56.9, a=1 ;area%:24.9, a=2;area%:9.6, a=;area%:3.7, a=4;area%:1.3

Synthetic Example 5

[0030] In a mixed solution of 10.0g of 48% of aqueous solution of sodium hydroxide and 155g of N,N'-dimethylace-toamide, 30.0g (0.12 mole) of BPS is added. After temperature is risen to 80°C and BPS is dissolved, 10.5g (0.06 mole) of α , α '-dichloro-p-xylene dissolved in 15g of xylene is dropped slowly. Then, ripened for 2 hours by same temperature. After ripened, the solution is poured into 900 ml of water and the crystallized solid is filtrated. The obtained crude crystalline is rinsed by methanol, filtrated and dried up, and 19.7g of white crystalline is obtained. Analyzed by

high precision liquid chromatograph, and the main components are identified as follows.

- α. α'-bis[4-(4-hydroxyphenylsulfonyl)phenoxy]-p-xylene 59.1%
- 4,4'-bis[4-(4-hydroxyphenylsulfonyl)phenyl-1,4-phenylenebismethyleneoxy] diphenylsulfone 23.1%.
- α , α' -bis[4-[4-(4-hydroxyphenylsulfonyl]phenyl-1,4-phenylenebismethylene oxy] phenylsulfonyl]phenoxyl-p-xylene 11.1%

[0031] In the present invention, when diphenylsulfone bridgeable type compound represented by general formula (1) or 3-{[(phenylamine)carbonyl]amino} benzenesulfoneamide compound represented by general formula (2) are used together with as the color developer for a higher temperature color developing layer, the mixing ratio is voluntarily decided along with the desired quality and not restricted. For instance, in a case in which the image preservability of higher temperature color developed image to plasticiser is especially important, if the amount of 3-{[(phenylamine) carbonyl]amino}benzenesulfoneamide is too small, the improving effect of image stability is not sufficient, on the contrary, if it is too much, said effect is deteriorated. By containing 0.1 to 0.5 parts of 3-{[(phenylamine)carbonyl] amino} benzenesulfoneamide to 1 part of diphenylsulfone bridgeable type compound a thermally sensitive recording medium whose recording sensitivity and image preservability are well balanced. When only 3-{[(phenylamine)carbonyl]amino} benzene sulfoneamide is used as the color developer for the higher temperature color developing layer, the color developed at lower temperature and the color developed at higher temperature can be clearly separated even if under the high temperature atmosphere.

[0032] In the present invention, as a color developer contained in the lower temperature color developing layer, for instance, bisphenol A type disclosed in Japanese Patent Laid Open Publication 3-207688 and Japanese Patent Laid Open Publication 5-24366, 4-hydroxybenzoic acid ester type, 4-hydroxyphtalic acid diester type, phtalic acid monoester type, bis-(hydroxyphenyl)sulfide type, 4-hydroxyphenylarylsulfone type, 4-hydroxyphenylarylsulfone type, 4-hydroxyphenylarylsulfone type, 4-hydroxybenzoyloxybenzoic acid ester type, bisphenol sulfone type, derivatives of aminobenzenesulfoneamide disclosed in Japanese Patent Laid Open Publication 8-59603 and diphenylsulfone bridgeable type compound disclosed in WO97/16420 can be mentioned. These mentioned color developers can be voluntarily chosen according to the desired recording sensitivity and other properties.

[0033] The typical examples of well-known color developer are mentioned below, however, not limited to them.

<Bisphenol A type>

[0034]

10

15

20

25

30

35

45

50

- 4.4'-isopropylidenediphenol (another name is bisphenol A)
- 4,4'-cyclohexylidenediphenol
- p,p'-(1-methyl-normalhexylidene)diphenol
- 1,7-di(hydroxyphenylthio)-3,5-dioxaheptane
- 40 <4-hydroxybenzoic acid ester type>

[0035]

- 4-hydroxybenzyl benzoate
- 4-hydroxyethyl benzoate
- 4-hydroxypropyl benzoate
- 4-hydroxyisopropyl benzoate
- 4-hydroxybutyl benzoate
- 4-hydroxyisobutyl benzoate
- 4-hydroxymethylbenzyl benzoate.
 - <4-hydroxyphthalic acid diester type>

[0036]

- 4-hydroxydimethylphthalate
- 4-hydroxydiisopropylphthalate
- 4-hydroxydibenzylphthalate

4-hydroxydihexylphthalate

<Phthalic acid monoester type>

[0037]

10

15

monobenzyl phthalate monocyclohexyl phthalate monophenyl phthalate monomethylphenyl phthalate monoethylphenyl phthalate monopropylbenzyl phthalate monohalogenbenzyl phthalate monoethoxybenzyl phthalate

<Bis-(hydroxyphenyl)sulfide type>

[0038]

bis-(4-hydroxy-3-tert-butyl-6-methylphenyl)sulfide 20 bis-(4-hydroxy-2,5-dimethylphenyl)sulfide bis-(4-hydroxy-2-methyl-5-ethylphenyl)sulfide bis-(4-hydroxy-2-methyl-5-isopropylphenyl) sulfide bis-(4-hydroxy-2,3-dimethylphenyl)sulfide bis-(4-hydroxy-2,5-dimethylphenyl)sulfide 25 bis-(4-hydroxy-2,5-diisopropylphenyl)sulfide bis-(4-hydroxy-2,3,6-trimethylphenyl)sulfide bis-(2,4,5-trihydroxyphenyl)sulfide bis-(4-hydroxy-2-cyclohexyl-5-methylphenyl)sulfide bis-(2,3,4-trihydroxyphenyl)sulfide 30 bis-(4.5-dihydroxy-2-tert-butylphenyl)sulfide bis-(4-hydroxy-2,5-diphenylphenyl)sulfide bis-(4-hydroxy-2-tert-octyl-5-methylphenyl)sulfide

35 <4-hydroxyphenylarylsulfone type>

[0039]

4-hydroxy-4'-isopropoxydiphenylsulfone 4-hydroxy-4'-n-propoxydiphenylsulfone 4-hydroxy-4'-n-butyloxydiphenylsulfone

<4-hydroxyphenylarylsulfonate type>

45 [0040]

40

- 4-hydroxyphenylbenzenesulfonate
- 4-hydroxyphenyl-p-tolylsulfonate
- 4-hydroxyphenylmethylenesulfonate 4-hydroxyphenyl-p-chlorobenzenesulfonate
- 4-hydroxyphenyl-p-tert-butylbenzenesulfonate
- 4-hydroxyphonyl-p-isopropoxybenzenesulfonate
- 4-hydroxyphenyl-1'-naphthalenesulfonate
- 4-hydroxyphenyl-2'-naphthalenesulfonate

55

<1.3-di[2-(hydroxyphenyl)-2-propyl]benzene type>

[0041]

- 1,3-di[2-(4-hydroxyphenyl)-2-propyl]benzene
- 1,3-di[2-(4-hydroxy-3-alkylphenyl)-2-propyl]benzene
- 1,3-di[2-(2,4-dihydroxyphenyl)-2-propyl]benzene
- 1,3-di[2-(2-hydroxy-5-methylphenyl)-2-propyl]benzene
- 10 < Resorcinol type>
 - [0042] 1,3-dihydroxy-6(α , α -dimethylbenzyl)-benzene.
 - <4-hydroxybenzoyloxybenzoic acid ester type>

[0043]

15

20

25

30

40

45

50

55

- 4-hydroxybenzoyloxybenzyl benzoate
- 4-hydroxybenzoyloxymethyl benzoate
- 4-hydroxybenzoyloxyethyl benzoate
- 4-hydroxybenzoyloxypropyl benzoate
- 4-hydroxybenzoyloxybutyl benzoate
- 4-hydroxybenzoyloxyisopropyl benzoate
- 4-hydroxybenzoyloxytert-butyl benzoate
- 4-hydroxybenzoyloxyhexyl benzoate
- 4-hydroxybenzoyloxyoctyl benzóate
- 4-hydroxybenzoyloxynonyl benzoate
- 4-hydroxybenzoyloxycyclohexyl benzoate
- 4-hydroxybenzoyloxy β-phenethyl benzoate
- 4-hydroxybenzoyloxyphenyl benzoate
 - 4-hydroxybenzoyloxy a -naphthyl benzoate
 - 4-hydroxybenzoyloxy β-naphthyl benzoate
 - 4-hydroxybenzoyloxysec-butyl benzoate
- 35 <Bisphenolsulfone type (I)>

[0044]

- bis-(3-1-butyl-4-hydroxy-6-methylphenyl)sulfone
 - bis-(3-ethyl-4-hydroxyphenyl)sulfone
 - bis-(3-propyl-4-hydroxyphenyl)sulfone
 - bis-(3-methyl-4-hydroxyphenyl)sulfone
 - bis-(2-isopropyl-4-hydroxyphenyl)sulfone
 - bis-(2-ethyl-4-hydroxyphenyl)sulfone
 - bis-(3-chloro-4-hydroxyphenyl)sulfone
 - bis-(2,3-dimethyl-4-hydroxyphenyl)sulfone
 - bis-(2,5-dimethyl-4-hydroxyphenyl)sulfone
 - bis-(3-methoxy-4-hydroxyphenyl)sulfone
 - 4-hydroxyphenyl-2'-ethyl-4'-hydroxyphenylsulfone
 - 4-hydroxyphenyl-2'-isopropyl-4'-hydroxyphenylsulfone
 - 4-hydroxyphenyl-3'-isopropyl-4'-hydroxyphenylsulfone
 - 4-hydroxyphenyl-3'-sec-butyl-4'-hydroxyphenylsulfone
 - 3-chloro-4-hydroxyphenyl-3'-isopropyl-4'-hydroxyphenylsulfone
 - 2-hydroxy-5-t-butylphenyl-4'-hydroxyphenylsulione
 - 2-hydroxy-5-t-aminophenyl-4'-hydroxyphenylsulfone
 - 2-hydroxy-5-t-isopropylphenyl-4'-hydroxyphenylsulfone
 - 2-hydroxy-5-t-octylphenyl-4'-hydroxyphenylsulfone
 - 2-hydroxy-5-t-butylphenyl-3'-chloro-4'-hydroxyphenylsulfone

2-hydroxy-5-t-butylphenyl-3'-methyl-4'-hydroxyphenylsulfone 2-hydroxy-5-t-butylphenyl-3'-isopropyl-4'-hydroxyphenylsulfone 2-hydroxy-5-t-butylphenyl-2'-methyl-4'-hydroxyphenylsulfone

<Bisphenolsulfone type (II)>

[0045]

10

15

25

35

40

50

55

4,4'-sulfonyldiphenol 2,4'-sulfonyldiphenol

3,3'-dichloro-4,4'-sulfonyldiphenol

3,3'-dibromo-4,4'-sulfonyldiphenol

3.3',5,5'-tetrabromo-4,4'-sulfonyldiphenol

3,3'-diamino-4,4'-sulfonyldiphenol

<others>

[0046]

p-tert-butylphenol
 2,4-dihydroxybenzophenone
 novolac type phenolic resin
 4-hydroxyacetophenone

p-phenylphenol

benzyl-4-hydroxyphenylacetate

p-benzylphenol

4,4'-bis(p-tolylsulfonylaminocarbonilamino)diphenylmethane

4,4'-bis(phenylaminotiocarbonylamino)diphenylsufide

[0047] Additionally, as an image forming material, metallic chelete type color developing components composed of higher fatty acid metal complex salt and polyhibric hydroxyaromatic compound disclosed in Japanese Patent Laid Open Publication 10-258577 can be used. Still more, these chelete color developing components can be used alone or can be used together with said leuco dye and color developer.

In this invention, a conventional well known sensitizer can be blended to each color developing layer so as to adjust the recording sensitivity, in the limitation in which the desired effect of this invention is not prevented. As an example of the sensitizer,

stearic acid amide, palmitic acid amide,

methoxycarbonyl-N-benzamidestearate,

N-benzoyl stearic acid amide,

N-icosenic acid amide,

ethylene-bis-stearic acid amide,

behenic acid amide,

45 methylene-bis-stearic acid amide,

methylolamide,

N-methylolsteric acid amide.

dibenzyl terephthalate.

dimethyl terephthalate,

dioctyl terephthalate,

p-benzyloxybenzylbenzoate.

1-hydroxy-2-phenylnaphthoate.

dibenzyloxalate

di-p-methylbenzyloxalate.

di-p-chlorobenzyloxalate,

2-naphthylbenzylether,

m-tarphenyl,

p-benzylbiphenyl.

4-biphenyl-p-tolylether,

di(p-methoxyphenoxyethyl)ether.

1,2-di(3-methylphenoxy)ethane.

1,2-di(4-methylphenoxy)ethane.

1.2-di(4-methoxyphenoxy)ethane,

1,2-di(4-chlorophenoxy) ethane.

1.2-diphenoxyethane.

1-(4-methoxyphenoxy)-2-(2-methylphenoxy)ethane.

p-methylthiophenylbenzylether,

1,4-di(phenylthio)buthane,

p-acetotoluidide,

10

15

20

25

30

35

40

45

55

p-acetophenetidide.

N-acetoacetyl-p-toluidine.

di-(β-biphenylethoxy)benzene,

p-di(vinyloxyethoxy)benzene,

1-isopropylphenyl-2-phenylethane

1,2-bis(phenoxymethyl)benzene

p-toluenesulfonamide.

o-toluenesulfonamide,

di-p-tolylcarbonate and

phenyl- α -naphtylcarbonate

can be mentioned, however is not intended to be limited to these compounds. These sensitizer can be used alone or by mixing more than two kinds of them.

[0048] As a binder to be used in the present invention, full saponificated polyvinyl alcohol of 200-1900 polymerization degree, partial saponificated polyvinyl alcohol, denatured polyvinyl alcohol by carboxyl, denatured polyvinyl alcohol by amide denatured polyvinyl alcohol by sulfonic acid, denatured polyvinyl alcohol by butylal modified polyvinyl alcohol, derivatives of cellulose such as hydroxyethyl cellulose, methyl cellulose, ethyl cellulose, carboxymethyl cellulose and acetyl cellulose, copolymer of styrene-maleic anhydride, copolymer of styrene-butadiene, polyvinyl chloride, polyvinyl acetal, polyacrylicamide, polyacrylic acid ester, polyvinylbutylal, polystyrene or copolymer of them, polyamide resin, silicon resin, petroleum resin, terpene resin, ketone resin and cumarone resin can be illustrated. These macromolecule compounds can be applied by being dissolved into solvents such as water, alcohol, ketone, ester or hydrocarbon or by being dispersed in water or other medium under an emulsion state or a paste state and these forms of application can be used in combination according to the quality requirement.

[0049] As a filler which can be used in this invention, an inorganic or an organic filler such as silica, calcium carbonate, kaoline, calcined kaoline, diatomaceous earth, talc, titanium dioxide, zinc oxide, aluminum hydroxide, polystyrene resin, urea-formaldehyde resin, copolymer of styrene-methacrylic acid, copolymer of styrene-butadiene or hollow plastic pigments can be mentioned.

[0050] If it is necessary to obtain more vivid color tone to a higher temperature color developing layer, it is effective to reduce the blending ratio of filler in higher temperature color developing layer or not to blend it. The reason why is presumed as follows. That is, in a case of color adding type, the color developing compound (reacted product of leuco dye and color developer, namely color developed image) formed in a higher temperature color developing layer at high temperature heating does not stay only in the higher temperature color developing layer and is absorbed in filler contained in a lower temperature color developing layer, then mixed effectively with the color developing compound formed in the lower temperature color developing layer, thus the higher temperature developed color tone, which is a mixed color, is obtained.

[0051] Still more, a parting agent such as metallic salt of fatty acid, a slipping agent such as waxes, an ultraviolet ray absorbing agent such as benzophenone type or triazole type, a water proof agent such as glyoxal, a dispersing agent, a defoamer, an antioxidant and a fluorescent brightening agent can be used.

[0052] As a substrate of the thermally sensitive recording medium, paper, recycled paper, synthetic paper, plastic film, foamed plastic film, non-woven cloth and metal foil can be used, further a complex sheet that combines these materials can be used.

[0053] Further, for the purpose to improve the friction resistance, it is possible to prepare an over coating layer composed of high polymer on the surface of the thermally sensitive recording layer. Furthermore, for the purpose to improve the sensitivity, it is possible to prepare an under coating layer containing organic or inorganic fillers between a color developing layer and a substrate.

[0054] The amount of color developer and leuco dye, the kind and amount of other additives to be used to each thermally sensitive recording layers of this invention are decided according to the required quality and recording feature.

and not restricted. However, in general, it is preferable to use 0 to 4 parts of filler to 1 part of color developer, and the desirable amount of a binder is 5 to 25 % to the total amount of solid. Further, it is desirable to contain 0.1 to 2 parts of leuco dye to 1 part of color developer.

[0055] Still more, in a case to use leuco dye which develops multi tone color in a higher temperature color developing layer and to obtain a complex tone color by high temperature heating, it is effective to make the amount of dye in the higher temperature color developing layer 0.5 to 3 parts, desirably 0.6 to 2.0 parts to 1 part of dye contained in the lower temperature color developing layer. And in a case to use a dye which develops mono tone color in a higher temperature color developing layer and in a lower temperature color developing layer and the color tone developed at high temperature heating is complex tone color, the ratio of each mono tone color type dye of higher temperature color developing layer and lower temperature color developing layer can be voluntarily adjusted according to the desired color tone.

[0056] In the present invention, the coating amount of each color developing layers are not restricted, however, ordinary are adjusted in the region of 1.5 to 12 g/m² by dry weight. Still more, in the present invention, it is possible to prepare a middle layer between higher temperature color developing layer and lower temperature color developing layer. In the coating which forms said middle layer, a binder, a filler, a sensitizer, a defoamer, an antioxidant, an UV absorber and a fluorescent brightening agent can be voluntarily blended. The coating amount of a middle layer can be adjusted between the region of 1 to 10 g/m².

[0057] Said organic color developer, dye and other additives which are added when needs arises, and are ground to the fine particles smaller than several microns diameter by means of a pulverizer such as a ball mill, an attriter or a sand grinder, or by means of an adequate emulsifying apparatus, then a binder and other additives are added when needs arises, thus the coating is prepared. As the concrete examples of a coating method, a coating by hand, a sizing press coater method, a roll coater method, an air knife coater method, a blend coater method, a flow coater method, a comma direct method, a gravure direct method, a gravure reverse method and a reverse-roll coater method can be mentioned. Or, it is possible to dry up the medium after spraying or dissolving the coating.

EXAMPLE

10

20

25

30

35

40

cpreparation of thermally sensitive recording medium>

[0058] The thermally sensitive recording medium of this invention will be illustrated more concretely by Examples, however, not intended to be limited to them. In the Examples and Comparative Examples, parts and % indicates weight part and weight %.

Example 1

[0059] Example 1 is an example that uses in a lower temperature color developing layer, 4-hydroxy-4'-isopropoxy-diphenylsulfone as a color developer and 3,3-bis(p-dimethylaminophenyl)-6-dimethylaminophtalide (blue color developing leuco dye) as a leuco dye, and in a higher temperature color developing layer, the compound of synthetic example 1 as a color developer, 3-(N-ethyl-p-toluidino)-6-methyl-7-anilinofluoran (black color developing leuco dye) as a leuco dye.

[0060] Dispersion of color developer for a lower temperature color developing layer (A solution), dispersion of leuco dye which develops blue color (B solution), dispersion of color developer for a lower temperature color developing layer (C solution) and dispersion of leuco dye which develops black color (D solution) are separately ground in wet condition to average particle diameter of 1 μ m by a sand grinder.

17

45

50

A solution (dispersion of color developer)	•
4-hydroxy-4'-isopropoxydiphenylsulfone	6.0 parts
10% polyvinyl alcohol	18.8 parts
water	11.2 parts
B solution (dispersion of blue color leuco dye)	
3.3-bis(p-dimethylaminophenyl)-6-dimethylaminophtalide	1.0 parts
10% polyvinyl alcohol	2.3 parts
water	1.3 parts
C solution (dispersion of color developer)	
compound of synthetic Example 1	6.0 parts
10% polyvinyl alcohol	18.8 parts
water	11.2 parts
D solution (dispersion of black color leuco dye)	
3-(N-ethyl-p-toluidino)-6-methyl-7-anilinofluoran	1.0 parts
10% polyvinyl alcohol	2.3 parts
water	1.3 parts

[0061] Then above mentioned dispersions are mixed by following ratio, stirred and dispersions are prepared.

[preparation of a coating for a lower temperature color developing layer]	
A solution (dispersion of color developer)	36.0 parts
B solution (dispersion of blue color leuco dye)	13.8 parts
SiO ₂ pigment (25% dispersion; Mizucasil P-527 (product of Mizusawa Chemical)	40 parts
10% polyvinyl alcohol	20 parts
[preparation of a coating for a higher temperature color developing layer]	
C solution (dispersion of color developer)	36.0 parts
D solution (dispersion of black color leuco dye)	13.8 parts
SiO ₂ pigment (25% dispersion; Mizucasil P-527 (product of Mizusawa Chemical)	20 parts
10% polyvinyl alcohol	20 parts

[0062] The prepared coating for a higher temperature color developing layer is coated on one side of substrate paper of 50 g/m² paper so as the coating amount to be 5.0g/m² and dried, and then over said surface the coating prepared for a lower temperature color developing layer is coated so as the coating amount to be 4.0g/m². Then the obtained sheet is treated by a super calendar so as the smoothness to be 500 to 600 sec., thus the multi chromatic thermally sensitive recording medium is obtained.

Example 2

10

15

20

25

30

35

45

50

55

[0063] A thermally sensitive recording medium is prepared same as to the Example 1. However, at the preparation of a high temperature color developing layer, SiO₂ pigment is not added.

Example 3

[0064] A thermally sensitive recording medium is prepared same as to the Example 2. At the preparation of A solution, 2,4'-sulfonyldiphenol is used instead of 4-hydroxy-4'-isopropoxydiphenylsulfone.

Examples 4, 5

[0065] A thermally sensitive recording medium is prepared same as to the Example 2. At the preparation of B solution. 3-(4-diethlamino-2-ethoxyphenyl)-3-(1-ethyl-2-methylindol-3-yl)-4-azaphthalide (blue color developing leuco dye: Example 4), 3,3-bis(1-n-butyl-2-methylindol-3-yl)phthalide (red color developing leuco dye: Example 5) are used instead

of 3.3-bis(p-dimethylaminophenyl)-6-dimethylaminophtalide.

Example 6

[0066] A thermally sensitive recording medium is prepared same as to the Example 2. At the preparation of C solution the compound of Synthetic Example 2 is used instead of the compound of Synthetic Example 1.

Examples 7, 8

[0067] A thermally sensitive recording medium is prepared same as to the Example 2. At the preparation of D solution, 3-dibutylamino-7-(o-chloroanilino) fluoran (black color developing leuco dye; Example 7), 3-dibutylamino-6-methyl-7-anilinofluoran (black color developing leuco dye; Example 8) are used instead of 3-(N-ethyl-p-toluidino)-6-methyl-7-anilinofluoran.

15 Example 9

[0068] A thermally sensitive recording medium is prepared same as to the Example 2. At the preparation of C solution 3-{[(phenylamine)carbonyl]amino} benzenesulfoneamide is used instead of the compound of Synthetic Example 1.

20 Example 10

25

30

[0069] A thermally sensitive recording medium is prepared same as to the Example 2. At the preparation of a coating for high temperature color developing layer, 3 parts of 3-{[(phenylamine)carbonyl]amino}benzenesulfoneamide dispersion (E solution) is added.

E solution (dispersion of color developer)	
3-{ [(phenylamine)carbonyl]amino}benzenesulfoneamide 10% polyvinyl alcohol	6.0 parts 18.8 parts
water	11.2 parts

Examples 11, 12, 13

[0070] A thermally sensitive recording medium is prepared same as to the Example 10. At the preparation of a coating for high temperature color developing layer, the adding amount of E solution is changed to 6 parts (Example 11), 18 parts (Example 12) and 27 pats (Example 13).

Comparative Example 1

[0071] A thermally sensitive recording medium is prepared same as to the Example 7. At the preparation of C solution, 4.4'-sulfonyldiphenol(bisphenol S) is used instead of the compound of Synthetic Example 1.

<Method for evaluation>

- [0072] Thermal recording is carried out on the prepared thermally sensitive recording media using Thermal sensitive printer (thermal head of ROHM Co., Ltd. type KM2004-A3 is installed), which is a product of MARKPOINT Co., Ltd. Printed by 0.076mj/dot impressive energy for lower temperature color developing and printed by 0.219mj/dot impressive energy for higher temperature color developing. Image density of recorded part is measured by means of a Macbeth densitometer (RD-914). At the measurement by Macbeth densitometer, red filter is used for blue color image, green filter is used for red color image and amber filter is used for black color image and ground part color.
 - [0073] As the heat resistance test, prepared lower temperature color developed specimen is placed in dry high temperature atmosphere of 65°C for 24 hrs and the color tone of lower temperature color developed part (0.076 mj/dot) is inspected by naked eyes of the inspector, further degree of color developing of ground part is evaluated by Macbeth densitymeter.
- [0074] Resistance to plasticiser test is carried out as follows. A single sheet of polyvinylchloride wrap (HIGHWRAP KMA: Mitsui Toatsu Chemicals Co., Ltd.) is wound round with 1 ply on a paper tube, stuck thereon a specimen developed by higher temperature (0.219 mj/dot) so as the recorded part to be upper surface, further wound round with 3 plies of the polyvinylchloride wrap, allowed to stand at 40°C for 24 hours, and measured the Macbeth density of the recorded

part. The obtained results are summarized in Table 1 and Table 2.

Table 1

Example	0.076 mj/dot		0.219 mj/dot	
	color tone	O.D.(image)	color tone	O.D.(image)
1	blue	1.16(R)	bluish black	1.34(A)
2	blue	1.17(R)	black	1.35(A)
3	blue	1.17(R)	black	1.35(A)
4	blue	1.21(R)	black	1.38(A)
5	red	1.22(G)	black	1.32(A)
6	blue	1.16(R)	black	1.33(A)
7	blue	1.16(R)	black	1.32(A)
8	light navy blue	1.20(R)	black	1.35(A)
9	blue	1.10(R)	black	1.34(A)
10	blue	1.18(R)	black	1.35(A)
11	blue	1.20(R)	black	1.37(A)
12	blue	1.20(R)	black	1.37(A)
13	blue	1.21(R)	black	1.36(A)
Co. Exp.1	blue	1.09(R)	black	1.36(A)

Example	hea	plasticiser resistance		
	color tone	O.D.(ground part)	O.D.(image)	
1	dark blue	0.08(A)	0.76(A)	
2	dark blue	0.08(A)	0.78(A)	
3	dark blue	0.08(A)	0.74(A)	
4	dark blue	. 0.09(A)	0.85(A)	
5	dark red	0.07(A)	0.81 (A)	
6	dark blue	0.09(A)	0.71 (A)	
7	dark blue	0.06(A)	0.74(A)	
8	navy blue	0.08(A)	0.84(A)	
9	blue	0.06(A)	0.20(A)	
10	dark blue	0.08(A)	0.81 (A)	
11	dark blue	0.08(A)	1.00(A)	
12	dark blue	0.08(A)	0.91 (A) **	
13	dark blue	0.07(A)	0.70(A)	
Co. Exp.1	black	0.15(A)	0.02(A)	
(Remarks)	in Tables, marks	in parenthesis indicates	the type of filter of Mac	

<Evaluation results>

[0075] From the Examples 1 to 13 of the present invention, sufficient recording density and clear color tone are obtained both on the results by lower impressive energy (0.076 mj/dot) and by higher impressive energy (0.219 mj/ dot). After the heat resistance test, fogging of the lower energy printed part (fogging of black color) and color developing of the ground part are slight, and color separation is clearly maintained, therefore, these products obtained by said Examples can be practically used. Especially, Example 9 that uses 3-{[(phenylamine)carbonyl] amino}benzenesulfoneamide alone indicates excellent result on heat resistance. Further, Examples 1 to 8 that use diphenylsulfone bridgeable type compound represented by general formula (1) are superior in the preservability of color developed image by high temperature. In Examples 10 to 13 that use diphenylsulfone bridgeable type compound together with 3-{[(phenylamine)carbonyl]amino}benzenesulfoneamide indicate good preservability of color developed image by high temperature. Especially, Examples 11 and 12 in which 0.1 to 0.5 parts of 3-[[(phenylamine)carbonyl] amino}benzenesulfoneamide is used to 1 part of diphenylsulfone bridgeable type compound, indicate excellent preservability of color developed image by high temperature. On the contrary, by the Comparative Example 1 that uses a color developer not specified in the present invention, sufficient results are obtained from the view point of recording density and color separation, however, the image printed at lower energy can not maintain it's initial clear color tone under the high temperature atmosphere and changes to black color, furthermore, the color developing of ground part is remarkable, and inferior in heat resistance and preservability of color developed image by high temperature.

20 Effect of the invention

[0076] The thermally sensitive recording medium of the present invention indicates good recording density at every developed color tone and clear developed color tone can be obtained. And, even if exposed under high temperature condition, it shows very clear color separation. Still more, since the color developing of ground part is small, the preservability of color developed image by high temperature is excellent, the present invention is suited to the uses which require heat resistance and image preservability and can be said very useful for the practical use.

Claims

10

15

25

30

35

40

45

50

55

1. A recording medium comprising:

at least one first color developer comprising a dye precursor and a developer component that reacts at a first temperature with said precursor to develop a first color; and and at least one second color developer that develops a second color, different from the first color, at a second temperature, lower than the first temperature;

wherein the developer component comprises:

(a) a diphenylsulfone bridgeable type compound of formula (1).

$$(R_1)_m$$
 $(R_2)_n$ $(R_3)_p$ $(R_4)_q$ $(R_5)_r$ $(R_6)_t$ $(R_7)_m$ $(R_7$

wherein each of R_1 to R_6 , which may be the same or different, is a halogen atom, an alkyl group of carbon number 1 to 6 or an alkenyl group,

each of m, n, p, q, r and t, which may be the same or different, is zero or an integer from 1 to 4 and when at least 2, the appropriate of R_1 s to R_6 s may be the same or different,

a is zero or an integer from 1 to 10.

each of X and Y, which may the same or different, is an unsaturated linear or grafted hydrocarbon group of carbon number 1 to 12 optionally possessing an ether bond, or

$$-R \longrightarrow R \longrightarrow (a) \qquad -CH_2 - \stackrel{T}{C} - CH_2 \longrightarrow (b)$$
or OH

wherein R is a methylene group or an ethylene group. T is a hydrogen atom or an alkyl group of carbon number 1 to 4; and/or

(b) 3-{[(phenylamine)carbonyl]amino}benzenesulfone amide compound of formula (2),

- 2. A medium according to claim 1, in which the developer component comprises (a) and (b).
- 3. A medium according to claim 1 or 2, which additionally comprises a support.

10

15

20

25

30

40

45

50

55

- 4. A medium according to claim 1, 2 or 3, in which the first and/or second color developers are in the form of layers.
- 5. A medium according to claim 4, in which the first and second color developer layers are laminated directly or indirectly to a support.
- 6. A medium according to claim 5, in which the first color developer layer is between the support and the second color developer layer.



Europäisches Patentamt

European Patent Office

Office européen des brevets



) EP 1 092 551 A3

(12)

EUROPEAN PATENT APPLICATION

(88) Date of publication A3: 08.05.2002 Bulletin 2002/19

(51) Int CI.7: B41M 5/30

(43) Date of publication A2: 18.04.2001 Bulletin 2001/16

(21) Application number: 00309043.8

(22) Date of filing: 13.10.2000

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU

MC NL PT SE

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 15.10.1999 JP 29394799 13.12.1999 JP 35315999

(71) Applicant: NIPPON PAPER INDUSTRIES CO., LTD. Kita-ku, Tokyo 114-0002 (JP)

(72) Inventors:

Hamada, Kaoru, c/o Res. Lab.,
 Product Development
 1-chome, Shinjuku-ku, Tokyo 161-0034 (JP)

Tadakazu, Fukuchi, Res. Lab.,
 Product Developm.
 1-chome, Shinjuku-ku, Tokyo 161-0034 (JP)

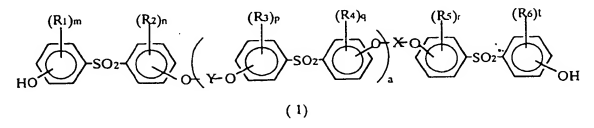
Midorikawa, Yoshimi c/o Res. Lab.,
 Product Develop
 1-chome, Shinjuku-ku, Tokyo 161.0034 (JP)

Kimura, Yoshihide, c/o Res. Lab.,
 Product Develop.
 1-chome, Shinjuku-ku, Tokyo 161-0034 (JP)

(74) Representative: McCluskie, Gail Wilson
 J.A. Kemp & Co.,
 14 South Square,
 Gray's Inn
 London WC1R 5JJ (GB)

(54) Multi chromatic thermally sensitive recording medium

(57) A multi chromatic thermally sensitive recording medium comprising, a higher temperature color developing layer containing dye precursor and an organic color developer that reacts with said dye precursor and develops color by heating, and at least one lower temperature color developing layer that develops different color from that of said higher temperature color developing layer and develops color by lower temperature than the color developing temperature of said higher temperature color developing layer laminated in order on a substrate, wherein an organic color developer contained in said higher temperature color developing layer contains said diphenylsulfone bridgeable type compound represented by general formula (1) and/or said 3-{(phenylamine)carbonyl] amino}benzenesulfoneamide compound represented by general formula (2),



wherein. X and Y can be different and indicates a saturate or an unsaturated liner or grafted hydrocarbon group of carbon number 1 to 12 which can possess an ether bond, or

$$-R$$
 (a)

or,

$$-CH2 - C-CH2 - (b)$$

wherein R indicates a methylene group or an ethylene group, T indicates a hydrogen atom or an alkyl group of carbon number 1 to 4,

and R_1 to R_6 independently indicate a halogen atom, an alkyl group of carbon number 1 to 6 or an alkenyl group, further m, n, p, q, r, t indicate an integer number of 0 to 4 and when are bigger than 2, R_1 to R_6 can be different and a is an integer of 0 to 10.



EUROPEAN SEARCH REPORT

Application Number EP 00 30 9043

ategory	DOCUMENTS CONSIDER Citation of document with incli		Relevant	CLASSIFICATION OF THE	
ategory	of relevant passag	es	to claim	APPLICATION (IntCl.7)	
A	PATENT ABSTRACTS OF vol. 017, no. 168 (M-31 March 1993 (1993-04 JP 04 329186 A (KALLTD), 17 November 1994 abstract *	-1391), D3-31) NZAKI PAPER MFG CO	1-6	B41M5/30	
A	EP 0 769 391 A (JUJO 23 April 1997 (1997-0 * claim l * * page 3, line 30 *	PAPER CO LTD) 04-23)	1-6		
A,D	EP 0 860 429 A (NIPPO 26 August 1998 (1998 * claim 1 * * page 3, line 24 ~	-08-26)	1-6		
A	US 3 895 173 A (ADACI 15 July 1975 (1975-0 * claim 1 *	HI KINICHI) 7-15)	1-6		
	-			TECHNICAL FIELDS SEARCHED (Int.CL7)	
				B41M	
	The present search report has be	een drawn up for all claims	7		
	Place of search	Date of completion of the search		Examinor	
	THE HAGUE	19 March 2002	Mai	rtins Lopes, L	
X : par Y : par doc A : led O : no	CATEGORY OF CITED DOCUMENTS ticularly relevant if taken alone ticularly relevant if tombined with anothe ticularly relevant if combined with anothe ticularly relevant if combined with anothe ticularly relevant hnological backgroundwritten discloture granddisc document	E : eartier pelent of after the filling to document cite L : document cite	T: theory or principle underlying the invention E: earlier patent document, but published on, or after the filling date D: document cled in the application L: document cled for other reasons 8: member of the same patent family, corresponding document		



Application Number

EP 00 30 9043

CLAIMS INCURRING FEES
The present European patent application comprised at the time of filling more than ten claims.
Only part of the claims have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims and for those claims for which claims fees have been paid, namely claim(s):
No claims tees have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims.
LACK OF UNITY OF INVENTION
The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:
see sheet B
All further search fees have been paid within the fixed time limit. The present European search report has been drawn up for all claims.
As all searchable claims could be searched without effort justifying an additional fee, the Search Division did not invite payment of any additional fee.
Only part of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid, namely claims:
None of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims, namely claims:

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 00 30 9043

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

19-03-2002

JP							
	04329186	A	17-11-1992	JP	2946811	B2	06-09-199
EP	0769391	A	23-04-1997	CA	2185846	A1	20-03-199
				DE	69601024	D1	07-01-1999
				DE	69601024	T2	15-07-1999
				EP	0769391	A1	23-04-199
				HK	1000908	A1	17-03-200
			•	JP	3029014	B2	04-04-200
				JP	9142034	A	03-06-199
				US	5753586	A	19-05-199
EP	0860429	A	26-08-1998	AU	707867	B2	22-07-199
				AU	7336496	A	22-05-199
				BR	9611435		23-03-199
			•	DE	69617991		24-01-200
				EP	0860429		26-08-199
				US	6037308		14-03-200
				CN	1200727		02-12-199
				WO	9716420		09-05-199
				JP 	10029969	A 	03-02-199
U\$	3895173	A	15-07-1975	JP	846476	7	28-02-197
				JP	48086543		15-11-197
				JP	51019989	-	22-06-197
			ह, प ्	JP	49034842		30-03-197
				JP	846488		28-02-197
				JP	49065240		25-06-197
				JP	51019990	-	22-06-197
				JP	846489	-	28-02-197
				JP	49065241		25-06-197
				JP	51019991	B	22-06-197 28-02-197
				JP	846490 49065242	-	25-06-197
				JP		_	22-06-197
				JP	51019992	_	24-04-197
				AU	460361 5214573		19-09-197
				AU	987103		13-04-197
				CA DE	2307774		23-08-197
				FR	2172386	_	28-09-197
				GB	1409831		15 -10- 197
				GB NL	7302182		21-08-197
					7 302162		21-00-197